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54) Title: DRUG TRIAL ASSAY SYSTEM		
(57) Abstract		

The invention provides a method for improving the efficacy of drug trials, the method comprising the step of screening samples from potential participants for the genetic basis of Gilbert's Syndrome and eliminating or including potential participants in a drug trial in the knowledge of them possessing or not possessing the genetic basis of Gilbert's Syndrome.

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"Drug Trial Assay System" 1 The present invention relates to drug trials, usually 3 carried out for or on behalf of pharmaceutical 4 companies. More particularly the invention relates to 5 a method for improving the efficacy of drug trials. 6 7 8 In the different stages of drug trials, regulatory authorities in different European countries and the FDA 9 in the USA require extensive data to be provided in 10 order to approve use of the drugs. 11 12 It is important that as much information as possible is 13 available in relation to all participants who take part 14 in drug trials, from volunteers who take part in phase 15 1 trials to patients involved in stage 3 clinical 16 17 trials. 18 In particular, if certain individuals or groups of 19 individuals have severe or abnormal reactions to drug 20 administration, further studies involving that drug 21 will be in jeopardy unless the reason for the reaction 22 23 is realised. 24 The knowledge of pharmacogenetics can play an important 25

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1 role in understanding the impact of drug metabolism on 2 pharmacokinetics, role of receptor variants in drug response and in the selection of patient populations 3 for clinical studies. 4 5 Considerable effort has been expended in attempting to 6 identify the pharmacogenetic basis of idiosyncsatic 7 adverse drug reactions, particularly hypersensitivity 8 9 reactions. While there is clear evidence for 10 pharmacogenetic influence on susceptibility to 11 hypersensitivity reactions, necessary and sufficient pharamacogenetic defects have not been identified. 12 13 14 The clinical implications of genetic polymorphism in 15 drug metabolism have been studied extensively (See 16 Tucker GT (1994) Journal Pharamacology 46 pages 417-17 424). 18 19 Gilbert's Syndrome (GS) is a benign unconjugated 20 hyperbilirubinaemia occurring in the absence of structural liver disease and overt haemolysis and 21 characterized by episodes of mild intermittent 22 23 jaundice. It is part of a spectrum of familial 24 unconjugated hyperbilirubinaemias including the more severe Crigler-Najjar (CN) syndromes (types 1 and 2). 25 26 GS is the most common inherited disorder of hepatic 27 bilirubin metabolism occurring in 2-12% of the 28 population and is often detected in adulthood through 29 routine screening blood tests or the fasting associated 30 with surgery/intercurrent illness which unmasks the hyperbilirubinaemia1.3. The most consistent feature in 31 32 GS is a deficiency in bilirubin glucuronidation but altered metabolism of drugs has also been reported3-5. 33 34 Altered rates of bilirubin production, hepatic haem production and altered hepatic uptake of bilirubin have 35 36 been reported in some GS patients2.

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Due to the benign nature of the syndrome and its 1 prevalence in the population it may be more appropriate 2 to consider GS as a normal genetic variant2 exhibiting a 3 reduced bilirubin glucuronidation capacity (which in 4 certain situations such as fasting, illness or 5 administration of drugs) could precipitate jaundice. 6 7 In drug trials where high levels of serum total 8 bilirubin is detected for certain individuals, it is 9 not clear whether this is because the individuals have 10 Gilbert's Syndrome or if it because of an effect of the 11 Whereas presently, results are explained merely 12 by saying that the individuals have Gilbert's Syndrome, 13 it is suspected that in the future, it will be 14 necessary to prove this fact. 15 16 Where a jaundiced phenotype is apparent after 17 volunteers have been accepted for a trial and have been 18 subjected to five days of a strict diet, no alcohol and 19 no smoking, the jaundiced appearance giving an 20 indication that the individuals have Gilbert's 21 Syndrome, may cause them to be ruled out of the trials. 22 Therefore, where approximately 250 individuals would be 23 required for phase 1 trials and about 6000 patients for 24 phase 3 trials, unnecessary time and effort would have 25 been spent during the first 5 days of these trials and 26 individuals having Gilbert's Syndrome may be ill 27 effected. 28 29 The present invention aims to provide a method of 30 improving the efficacy of drug trials in view of the 31 problems mentioned above. 32 33 According to the present invention there is provided a 34 method for improving the efficacy of drug trials, the 35 method comprising the step of screening samples from 36

4

1 individuals for the genetic basis of Gilbert's 2 Syndrome. 3 In a prefered embodiment of the invention the method 4 comprises the steps taking a sample from each potential 5 participant in a drug trial, screeing the samples for 6 7 the genetic basis of Gilbert's Syndrome, identifying 8 participants having the genetic basis of Gilbert's 9 Syndrome. 10 11 The sample may comprise blood, a buccal smear or any 12 other sample containing DNA from the individual to be tested. 13 14 In one embodiment the method comprises the further step 15 of eliminating participants having the genetic basis of 16 17 Gilbert's Syndrome from the drug trial. 18 In an alternative embodiment, the method can comprise 19 the further step of selecting participants having the 20 genetic basis of Gilbert's syndrome and eliminating 21 others from the drug trial. 22 23 In a further alternative the results of the drug trials 24 can be interpreted in the knowledge that certain 25 participants have Gilbert's Syndrome. 26 27 Preferably the method comprises the steps of isolating 28 DNA from each sample, amplifying the DNA in a region 29 indicating the genetic basis of Gilbert's Syndrome, 30 isolating amplified DNA fragments by gel 31 32 electrophoresis and identifying individuals having the genetic basis of Gilbert's disease. 33 34 Preferably the DNA is amplified using the polymerase 35

chain reaction (PCR) using a radioactively labelled

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pair of nucleotide primers.
  1
  2
       The primers are designed to prime the amplification
  3
       reaction at either side of an area of the genome known
  4
 5
       to be associated with Gilbert's Syndrome.
  6
 7
       Preferably the DNA region indicating the genetic basis
 8
       of Gilbert's Syndrome is the gene encoding UDP-
 9
       glucuronosyltransferase (UGT).
10
11
      By gene is meant, the non coding and coding regions and
12
      the upstream and downstream noncoding regions.
13
14
      In a preferred embodiment the DNA to be amplified is in
15
      an upstream promoter region of the UGT1*1 exon1.
16
17
      Most preferably the DNA to be amplified includes the
18
      region between -35 and -55 nucleotides at the 5' end of
19
20
      UGT1*1 exon.
21
      According to the invention there are provided suitable
22
      primers for use in a PCR reaction including primer
23
24
    . pairs;
25
26
      A/B(A,5'-AAGTGAACTCCCTGCTACCTT-3',
27
      B,5'-CCACTGGGATCAACAGTATCT-3') or
28
      C/D (C,5'-GTCACGTGACACAGTCAAAC-3';
      D 5'-TTTGCTCCTGCCAGAGGTT-3')
29
30
      The invention further comprises a kit for screeing
31
      individuals for participation in drug trials, the kit
32
      comprising primers for amplifying DNA in a region of
33
      the genome indicating the genetic basis of Gilbert's
34
35
      Syndrome.
36
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1 Using primer sequences as described herein, DNA can be 2 amplified and analysed using among others any of the 3 following protocols; 5 Protocol 1 Radioactive method 6 7 1. Extract DNA from Buccal Cells or 3ml Blood. 8 9 10 2. Choose primers from either side of the "TATA" box 11 region of UGT1*1 exon1 regulatory sequence. 12 Freshly end label one primer with $[\gamma^{32}\alpha]$ -ATP (40 13 min). 14 15 3. Amplifying a small region up to 100 bp in length 16 by PCR (2h). 17 18 4. Apply to 6% PAG denaturing gel (preparation, 19 loading, run time, 4h). 20 21 5. Expose (-70°C) wet gel to autoradiographic film 22 (15 min). 23 24 This method takes about 7h to complete. Polymorphisms 25 only observed in TATA box non coding region todate. 26 27 Protocol 2 28 Alternative Radioactive Method: Solid Phase 29 Minisequencing 30 31 Extract DNA (as above) 1. 32 33 2. Prepare primers biotinylating one 34 35 Amplify DNA by PCR using primers 3.

7

Captive biotinylated PCR products on streptavidin 4. 1 coated support and deactive. 3 Carry out primer extension reaction sequencing. 4 5. 5 Protocol 3 6 7 Non-Radioactive Methods: 8 (a) Analysis by Single Strand Conformational 9 Polymorphism (SSCP) 10 Extract DNA (as above). 11 1. 12 13 2. Choose primers either side of the TATA Box. 14 15 З. Amplify a small region up to 100 bp in length by 16 PCR (2H). Denature and place on ice (15 min). 17 4. 18 19 5. Load onto a non-denaturing PAG gel, (preparation/load/run time, 4h). 20 21 Stain with Ethidium bromide or silver nitrate (30 22 6. 23 mm). 24 25 This method still takes about 7h to complete, but is 26 potentially slightly cheaper since there is no 27 radioactivity or autoradiography. 28 29 This method could be done on an automated DNA sequencer from stage 5, if primers are tagged with chromophores 30 in PCR stages 2 and 3. Result would then be read 31 automatically. 32 33 34 (b) Oligonucleotide Assay Hybridization 35

36

1.

Extract DNA (as above).

8

Choose primers and amplify DNA by PCR up to 100 bp
 in length.

3

4 3. Apply DNA to plastic grids.

5

4. Screen bound DNA samples with specific DNA probes
 for TA₅, TA₆, TA₇ tagged with different
 coloured/fluorescent chromphores.

9

10 5. Read ouput automatically for experimental protocols.

12

13 References

14

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16

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The basis of the invention is illustrated in the 1 following example with reference to the accompanying 2 figures wherein: 3 Figure 1 illustrates genotypes at the TATA box sequence 5 upstream of the UGT1*1 exon 1 determined by direct 6 sequencing and radioactive PCR. 7 8 Figure 2 illustrates serum total bilirubin (μ mol/1) 9 10 plotted against UGT1*1 exon 1 genotype. 11 12 Figure 3 illustrates segregation of the 7/7 genotype 13 with elevated serum total bilirubin concentration in a family with GS. 14 15 16 Figure 4 illustrates the 5' sequence of the UGT1*1 exon 17 1 and the position of the primers with respect to the 18 UGT gene. 19 20 Example 21 22 We have examined the variation in the serum total 23 bilirubin (STB) concentration in a representative group 24 of the Eastern Scottish population (drug-free, alcohol-25 free non-smokers) in relation to genotype at the UDPglucuronosyltransferase subfamily 1 (UGT1) locus. 26 27 Subjects with the 77/7 genotype in this population have 28 a significantly higher STB than those with 6/7 or 6/6 29 genotypes. Of 14 control subjects who underwent a 24 30 hour fast to establish whether they had Gilbert Syndrome (GS), only 7/77 subjects had GS. In addition, 31 32 one confirmed GS patient, two recurrent jaundice 33 patients and 9 clinically diagnosed GS patients had the 34 7/7 genotype. Segregation of the 7/7 genotype with elevated STB concentration has also been demonstrated 35 in a family of 4 Gilbert members. This incidence of 36

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the 7/7 genotype in the population is 10-13%. Here, we 1 2 demonstrate a correlation between variation in the 3 human STB concentration and genotype at a TATA sequence upstream of the UGT1*1 exon 1 and that the 7/7 genotype 4 5 is diagnostic for GS. 6 7 The inheritance of GS has been described as autosomal dominant or autosomal dominant with incomplete 8 penetrance based on biochemical analysis6. More recent 9 10 reports have suggested that the mildly affected 11 (Gilbert) members of families in which CN type 2 (CN-2) 12 occurs are heterozygous for mutations in the UDI3glucuronosyltransferase subfamily 1 (UGT1) gene which 13 14 cause CN-2 in the homozygous state. The inheritance of 15 GS in these families is autosomal dominant while CN-2 16 is autosomal recessive ?-!!. However, the incidence of 17 CN-2 in the population is very rare and the frequency 18 of alleles causing CN-2 would not be sufficient to 19 explain the population incidence of GS. 20 An abstract by Bosma et al12 suggested a correlation 21 22 between homozygosity for a 2bp insertion in the TATA 23 box upstream of UGT1*1 exon 1 and GS (no mutations were 24 found in the coding sequence of the UGT1*1 gene). 25 this report we demonstrate that the primary genetic 26 factor contributing to the variation in the serum total 27 bilirubin (STB) concentration in the Eastern Scottish population is the sequence variation reported by Bosma 28 29 et al 12 . In addition, we show that the 7/77 genotype is associated with GS and occurs in 10-13% of the 30 31 population. 32 33 Methods 34 Patients and Controls 35 Whole blood (3ml) was collected into EDTA(K3) Vacutainer tubes (Becton Dickinson) from one confirmed 36

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male Gilbert patient (diagnosed following a 48 hour 1 restricted diet13), two female patients with recurrent 2 jaundice/associated elevated STB (29-42 μ mol/1) and 9 3 (1 female, 8 male) clinically diagnosed GS subjects (persistent elevation of the STB amidst normal liver 5 function tests.) The patients were aged 22-45 years. 6 7 77 non-smoking residents selected at random from the 8 Tayside/Fife region of Scotland (39 females aged 19-58 9 years, mean 32.41± 10.94; 38 males aged 23-57, means 10 35.58 ± 9.04) participated in this study. Whole blood 11 (9ml) was collected 8-10am) into EDTA(K3) Vacutainer 12 tubes (Becton Dickinson) for DNA extraction and SST 13 Vacutainer tubes (Becton Dickinson) for biochemical 14 investigations. The subjects had not taken any 15 medication or alcohol in the previous 5-7 days and had 16 fasted overnight (12 hours). 14 controls subsequently 17 underwent further biochemical tests (following a 3 day 18 abstinence from alcohol) before and after a 24 hour 19 400-calorie diet14 to determine if they had GS. 20 patients/controls were fully informed of the study and 21 gave consent for their blood to be used in this study. 22 23 Biochemistry and DNA Extraction 24 25 The following biochemical tests were performed on 26 control blood samples; alanine aminostransferase, 27 albumin, alkaline phosphatase, amylase, STB, 28 cholesterol, creatinine, creatine kinase, free 29 thyroxine, gamma-glutamyl-transferase, glucose, HDL-30 cholesterol, HDL-cholesterol/total cholesterol, iron, 31 lactate dehydrogenase, percentage of saturated 32 transferrin (PSAT), proteins, serum angiotensin 33 converting enzyme, thyroid stimulating hormone, 34 transferrin, triglycerides, urate, urea. 14 controls 35 also had pre- and post-fasting (24 hour) alanine 36

12

1 aminostransferase, albumin, alkaline phosphatase, STB 2 and urate measured. DNA was prepared using the Nucleon 3 II Genomic DNA Extraction Kit (Scotlab) according to 4 manufacturer's instructions. 5 6 Genotyping 7 8 Polymerase Chain Reaction 9 10 Primer pairs A/B (A, 5'-AAGTGAACTCCCTGCTACCTT-3'; B, 11 5'-CCACTGGGATCAACAGTATCT-3') or C/D (C,5'-12 GTCACGTGACACAGTCAAAC-3';D, 5'-TTTGCTCCTGCCAGAGGTT-3') 13 flanking the TATA box sequence upstream of the UGTI*1 14 exon 1 were used to amplify fragments of 253-255bp and 15 98-100bp, respectively. Amplifications (50µl) were 16 performed in 0.2mM of each deoxynucleoside triphosphate 17 (dATP, dCTP, dGTP, dTTP), 50mM KCI, 10mM Tris.HCl (pH 9.0 at 25°C), 0.1% Triton X-100, 1.5mM MgCl₂, 0.25 μ M of 18 19 each primer, 1 Unit of Tag Polymerase (Promega) and 20 human DNA $(0.25-0.5\mu g)$. The polymerase chain reaction 21 (PCR) conditions using the Perkin-Elmer Cetus DNA 22 Thermal Cycler were: 95°C 5 min followed by 30 cycles 23 of 95° 30 sec, 58°C 40 sec, 72°C40 sec. 24 25 Direct Sequencing 26 27 Amplification was confirmed prior to direct sequencing by agarose gel electrophoresis. Sequencing was 28 performed using $[\alpha^{-15}S]$ -dATP (NEN Dupont) with the USB 29 Sequenase™ PCR Product Sequencing Kit according to 30 31 manufacturer's instructions. Sequenced products were 32 resolved on 6% denaturing polyacrylamide gels. 33 dried gels were exposed overnight to autoradiographic film prior to developing. 34 35 36 Radioactive PCR

13

Amplification was performed as above using primer pair 1 C/D except that 2.5 pmol of primer C was radioactively 2 5' end-labelled with 2.5 μ Ci of [γ - 32 P]-ATP (NEN Dupont) 3 prior to amplification. Products were resolved on 6% denaturing polyacrylamide gels and the wet gels exposed 5 to autoradiographic film (-70°C 15 min) and the 6 autoradiographs developed. 7 Statistics 9 10 A t-test was used to determine if there was a 11 significant age difference between males and females. 12 χ^2 analysis was used to assess any difference in the 13 distribution of the 6/6, 6/7 and 7/7 genotypes in males 14 and females and also to determine if the 7/7 subjects 15 from the 24 hour fasted group had STB elevated into the 16 range diagnostic for GS14. An analysis of variance was 17 performed to compare mean STB in males and females 18 within each genotype group. A non-parametric test, the 19 Mann-Whitney U-Wilcoxon Rank Sum W Test was used to 20 determine whether there was a significant difference in 21 mean STB between males and females (irrespective of 22 genotype). Correlations and significance tests were 23 performed for STB versus PSAT and STB versus iron. A 24 probability (p) of (0.05 was accepted as significant. 25 26 27 Results 28 In Figure 1 a photographic representation of the sense 29 DNA sequences obtained by PCR/direct sequencing of DNA 30 samples having the genotypes 6/6, 6/7 and 7/7 is shown. 31 The common allele, (TA), TAA, is denoted by "6" while the 32 rarer allele, (TA), TAA, is denoted by "7". Below each 33 sequence is an overexposed photographic representation 34 of the 98 to 100bp resolved fragments amplified using 35 primer pair C/D which flank the TATA sequence upstream 36

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1 of the UGT1*1 exon 1. The additional fragments of 99 2 and 101 bases are thought to be artifacts of the PCR 3 process where there is non specified addition of an extra nucleotide to the 3' end of the amplified 4 product²¹. Figures 1b illustrates results after testing 5 6 a range of unknown individuals. 7 8 In Figure 2 males (M) and females (F) are plotted 9 separately. Each circle/square represents the result of a single control subject. The squares indicate the 10 14 controls who also underwent the 24 hour restricted 11 12 diet (see Methods). The filled circles/squares 13 represent those who had a lower than normal PSAT (≤ 22%) while the half-tone circles represent those who 14 had a higher than normal PSAT (≥ 55%). The mean STB 15 16 concentrations (indicated by the horizontal lines) for 17 males were $13.24 \pm 3.88 (6/6)$, $13.94 \pm 6.1 (6/7)$ 18 including control h or 12.69 ± 3.34 excluding control h, 29 \pm 14.45 (7/7) and for females were 9 \pm 3.62 19 (6/6), 12.2 ± 3.53 (6/7), 21.6 ± 7.8 7/7). 20 encircled result is from control h (discussed in the 21 22 text). 23 24 In Figure 3 males and females are represented by 25 squares and circles, respectively. Filled and halffilled circles/squares indicate the genotypes 7/7 and 26 27 6/7, respectively. The numbers in parentheses below 28 each member of the pedigree are the STB concentrations measured after a 15 hour fast and 7 day abstinence from 29 alcohol. All family members were non smokers who were 30 31 not taking any medication when the biochemical tests 32 were performed. Elevated STB are underlined. Individual members of each generation (I or II) are 33 denoted by the numbers 1-4 above each circle/square. 34 Generation III have not yet been tested. 35 36

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There was no significant age difference between males
 1
      and females (t = -1.38, p = 0.17). Genotypes were
 2
      determined initially by amplification/sequencing and
 3
      later by the radioactive PCR approach. Individuals
 4
      homozygous for the common allele, hetrozygous or
 5
      homozygous for the rarer allele have the genotypes 6/6,
 6
      6/7 and 7/7, respective. 12 DNA samples (2 of 6/6, 3
 7
      of 6/7 and 4 of 7/7) were analysed by both methods and
 8
      genotype results were identical (see Figure 1).
 9
10
      Genotype frequencies in male controls were 6/6 (44.74%,
11
      6/7 (44.74%), 7/7 (10.52%) and in female controls were
12
      6/6 (35.9%), 6/7 (51.3%), 7/7 (12.8%). There was no
13
      significant difference between the genotype proportions
14
      in the two groups (\chi^2 = 0.6 at 2 df, p = 0.7). Control
15
      h (encircled in Figure 2) had a STB which was 2.4 SD
16
      above the mean STB for that group (mean calculated
17
      including control h). The results for control h were
18
      repeatable and he is currently being investigated to
19
      exclude haemochromatosis. Comparison of mean STB in
20
      males and females revealed that females have a
21
      significantly lower concentration than males (p = 0.031
22
      including control h; p + 0.0458 excluding control h).
23
      There was a strong correlation between genotype and
24
      mean STB concentration within the control group (p (
25
      0.001) irrespective of whether control h was included
26
      and there was a significant difference in mean STB
27
      between males and females of the same genotype (p (
28
      0.05) irrespective of whether control h was included
29
      (see Figure 2). All patients studied had the 7/77
30
      genotype.
31
32
     Correlations between STB/PSAT (r = 0.4113, p =
33
     0.001) (see Figure 2) and STB/iron females (p = 0.001)
34
     than males (p = 0.01) but when control h is excluded
35
     there was no significant correlation in males.
36
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The STB concentrations of control who underwent the 24 2 hour restricted diet (see Methods) are shown in Table The normal fasting response is a small rise in the 3 base-line STB (not exceeding a final concentration of 25μmol/1) most of which is unconjugated while GS 5 patients have a lone biochemical feature a raised STB 6 ()25 μ mo1/1 but (50 μ mo1/1) most of which is 7 unconjugated14. The 6/6 and 6/7 controls had post-8 fasting STB of ≤23μmol/1 while all 7/77 controls were 9 ≥31µmo1/1. Other liver function tests were within 10 acceptable ranges for the age and sex of the subjects. 11 The 7/77 genotype correlates with a fasted STB (24 12 hour) within the range diagnostic for GS14 (p (13 0.01) (see Table 1). In addition, the 7/7 genotype 14 segregates with elevated STB concentration in a family 15 with 4 GS members (Figures 3). 16 17 18 Table 1 shows a comparison of the UGT1*1 exon 1 genotype with elevation in the serum total bilirubin 19 after a 24 hour 400-calorie restricted diet14. 20 21 22 An elevation of the fasting STB to a final concentration in the range $25-50\mu\text{mol/l}$ is considered to 23 be diagnostic for GS14. The 7/7 subject denoted by * 24 has a fasting and non-fasting STB of \rangle 50 μ mol/l but 25 this value is within a range considered by others to 26

conform to a diagnosis of GS7-11.

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Table 1

		24 hour fast		
Genotype	Sex	Before	After	Fasting bilirubin >25 & <50µmol/l
6/6	M M M	8 9 12	17 19 15	ио ио ио
6/7	F F F M M	8 9 11 12 8 15	17 13 12 17 10 23 18	NO NO NO NO NO NO
7/7	F F M	9 12 19 62	34 34 31 96	YES YES YES NO*

Discussion

A few recent reports claim to have identified the genetic cause of GS^{10-12} . Clinical diagnosis of GS is often based on a consistent midly elevated non-fasting STB ()17 μ mol/1) as the sole abnormal liver function test, intermittent jaundice or both. The diagnosis can be confirmed by elevation of the STB to 25-50 μ mol/1 after a 24 hour 400-calorie diet¹⁴ or by elevation of the unconjugated bilirubin by) 90% within 48 hours of commencing a 400 calorie diet¹³.

Sato's research group recently reported the occurrence of 7 different heteroxygous missence mutations in unrelated Gilbert patients (most of the mutations have been found in the homozygous state in affected members of CN families), however, the non-fasted STB for these patients were \rangle 52 μ mo1/1 (with the exception of one,

18

1 31μ mo1/1)^{10,12}. These non-fasted STB concentrations already exceed the diagnostic range for GS14, hence 2 3 these patients have a more severe form of hyperbilirubinaemia than those studied in this report, 4 while those in the Bosma et al 2 abstract had STB 5 concentrations similar to those studied here. 6 7 The example herein shows that the variation in the STB 8 9 levels after an overnight fast (and in the absence of exposure to known inducers of the UGT1*1 isoform in GS, 10 such as alcoholic15 and drugs16) a representative group 11 12 of the Eastern Scottish population is primarily due to (or associated with) the TATA box sequence variation 13 reported by Bosma et al¹². In agreement with previous 14 work females have a significantly lower mean STB 15 concentration than males 17-18. 16 17 Individuals with the 7/7 genotype in the population 18 have GS (see Table 1). One of the 7/7 controls 19 indicated in Table 1 had a non-fasting STB similar to 20 those reported for heterozygous carriers of CN-2 21 mutations⁷⁻¹¹ which suggests that this subject may also 22 be a carrier of a CN-2 mutation, alternatively, the 23 very elevated bilirubin in this patient may be due to 24 the coexistence of Reavon's Syndrome (characterized by 25 a collection of abnormal biochemical results which are 26 risk factors for coronary heart disease) 19. 27 28 We have found that 10-13% of the Eastern Scottish 29 population have the genotype associated with mild GS. 30 31 None of the Gilbert subjects from the control population were aware that they had an underlying 32 metabolic defect in glucuronidation with testifies to 33 its benign nature. Three 7/7 controls had STB 34 concentrations comparable to mean levels observed in 35 heterozygotes, however, they also had a lower than 36

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normal PSAT (≤22%) (see Figure 2). The observed

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correlation between STB and PSAT (p = 0.001) (Figure 2) 2 and STB and iron (females p = 0.001 and males p = 0.013 including control h) indicates that other genetic and 4 environmental factors affecting the serum PSAT and iron 5 values will in turn affect the STB concentration. 6 7 From the data presented here and previous reports it 8 seems clear that there are mild and more severe forms 9 The milder form (fasted STB $25-50\mu\text{mol/1}$) is 10 either caused by (or is associated with) a homozygous 11 2bp insertion at the TATA sequence upstream of the 12 UGT1*1 exon 1 (autosomal recessive inheritance) while 13 the rarer more severe dominantly inherited forms 14 identified to date $^{7.11}$ (non-fasted STB) 50 μ mol/l are due 15 to heterozygosity for a mutation in the coding region 16 of the UGT1*1 gene which in its homozygous state causes 17 The particular genetic abnormality causing GS in 18 a patient will have implications for genetic 19 counselling as the dominantly inherited form of two GS 20 patients could result in offspring with CN-2, whereas 21 the recessive form in one or both GS patients would 22 have less serious implications. It is important to 23 discriminate between the two forms and provide suitable 24 genetic counselling for such couples. The rapid DNA 25 test presented here (less than 1 day for extracted DNA) 26 carried out in addition to biochemical tests following 27 a 12 hour overnight fast (without prior alcohol or drug 28 intake would permit such a diagnosis. The compliance 29 rate for the current 24 and 48 hour restricted diet 30 tests for GS13-14 is debatable and hence the overnight 31 fast has obvious advantages and only one blood sample 32 or a buccal smear is required (for genetic and 33 biochemical analysis) in contrast to the 2-3 blood 34 samplings required for the 24 and 48 hour tests. 35 approach to GS testing would be cost effective in terms 36

20

1 of fewer patient return visits to clinics and in 2 identifying couples at risk of having children with 3 CN-2. 4 In addition, the recent finding of an increased 5 6 bioactivation of acetominophen (a commonly used 7 analgesic which is eliminated primarily by glucuronidation) in GS patients indicates the greater 8 9 potential for drug toxicity in these patients if 10 administered drugs which are also conjugated by UGT1 isoforms3. In fact, ethinylestradiol (EE2) has recently 11 12 been shown to be primarily glucuronidated by the UGT1*1 isoform in man²⁰ and hence this could have implications 13 14 for female Gilbert patients taking the oral 15 contraceptive who are then more predisposed to developing jaundice. 16 17 18 19 The tests outlined herein have obvious implications for setting up drug trials in understanding unusual results 20 in ruling out individuals who may be adversely affected 21 22 by the drugs or in positively choosing these individuals to determine the effects of particular 23 drugs on hyperbilirubinaemia. 24 25

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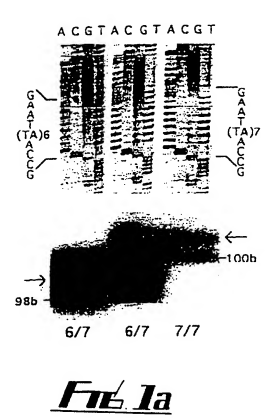
1	CLA	IMS
2		
3	1.	A method for improving the efficacy of drug
4		trials, the method comprising the step of
5		screening samples from potential participants for
6		the genetic basis of Gilbert's Syndrome and
7		eliminating or including potential participants in
8		a drug trial in the knowledge of them possessing
9		or not possessing the genetic basis of Gilbert's
10		Syndrome.
11		
12	2.	A method as claimed in claim 1 comprising the
13		steps of:
14		
15		 a) taking a sample from each potential
16		participant in a drug trial,
17		
18		b) screening the samples for the genetic basis
19		of Gilbert's Syndrome,
20		
21		 c) identifying participants having the genetic
22		basis of Gilbert's Syndrome, and
23		
24		d) proceeding with drugs trials in the knowledge
25		of participants possessing or not possessing
26		the genetic basis of Gilbert's Syndrome.
27		
28	3	A method as claimed in claim 1 or 2 wherein the
29		sample is chosen from blood, buccal smear or any
30		other sample containing DNA from the potential
31		participants.
32		
33	4.	A method as claimed in any of the preceding claims
34		further comprising the step of eliminating
35		participants having the genetic basis of Gilbert's
36		Syndrome from a drugs trial.

,	5	A method as claimed in any of claims 1 to 3
1	5.	wherein the method comprises the further step of
2		
3		selecting only participants having genetic basis
4		for Gilbert's Syndrome for a drugs trial.
5		
6	6.	A method as claimed in any of claims 1 to 3
7		further comprising the step of interpreting the
8		results of the drugs trial in the knowledge that
9		certain participants have Gilbert's Syndrome.
10		
11	7.	A method as claimed in any of the preceding claims
12		wherein the method comprises the steps of:
13		
14		 a) isolating DNA from each sample,
15		
16		 b) amplifying the DNA inner region indicating
17		the genetic basis for Gilbert's Syndrome,
18		
19		 c) isolating amplified DNA fragments, and
20		
21		d) identifying individuals having the genetic
22		basis of Gilbert's Syndrome.
23		
24	8.	A method as claimed in any of the preceding claims
25		wherein the DNA is amplified using the polymerase
26		chain reaction (PCR) using a radioactively
27		labelled pair of nucleotide primers.
28		Table of March 1997
29	10.	A method as claimed in any of claims 7 to 9
30	10.	wherein the DNA region indicating the genetic
31		basis of Gilbert's Syndrome is the gene encoding
32		UDP-glucuronosyltransferase (UGT).
		ODF-GIUCUIONICIANSIEIANSE (OGI).
33		A method on albimed in the of albims 7 to 10
34	11.	A method as claimed in any of claims 7 to 10
35		wherein the DNA to be amplified is in an upstream
36		promoter region of the UGT 1*1 exon 1.

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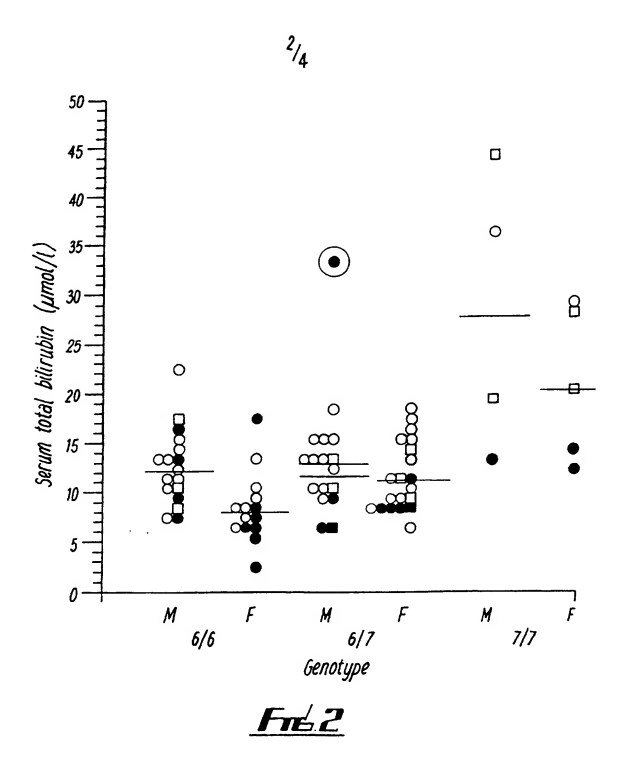
1	12.	A method as claimed in any of claims 7 to 11
2		wherein the DNA to be amplified includes the
3		regions between -35 and -55 nucleotides at the 5'
4		end of UGT 1*1 exon.
5		
6	13.	A kit for screening individuals participation in
7		drug trials, the kit comprising primers for
8		amplifying DNA in the region of the genome
9		indicating the genetic basis of Gilbert's
10		Syndrome.
11		
12	14.	Primers for use in a method as claimed in any of
13		the preceding claims including primer pairs, AB or
14		CD as follows:
15		
16		A/B(A,5'-AAGTGAACTCCCTGCTACCTT-3',
17		B,5'-CCACTGGGATCAACAGTATCT-3') or
18		C/D (C,5'-GTCACGTGACACAGTCAAAC-3';
19		D 5'-TTTGCTCCTGCCAGAGGTT-3').

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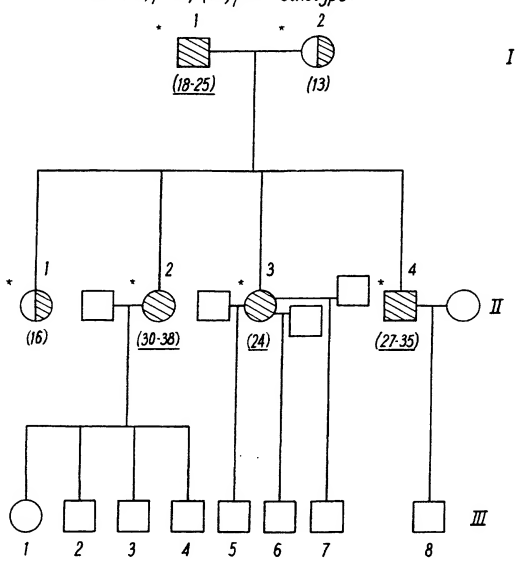




Fre 1b



Pedigree Showing Segregation of the Gilbert Phenotype with 3/4 the (TA), TAA / (TA), TAA Genotype



I, II, III - generations in family * = genetic and biochemical data available

☐ male

☐ ○ homozygotes for the (TA)7 TAA allele

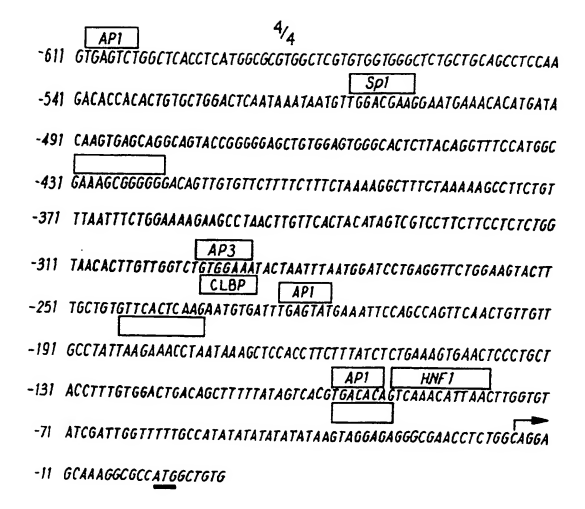
O female

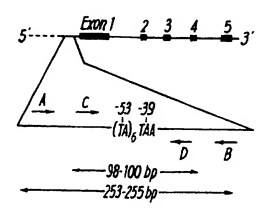
heterozygotes for the (TA), TAA and (TA), TAA alleles

(13) = total serum bilirubin

(18-25) = elevated total serum bilirubin







Fil. 4

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